# 24,25,28-Trihydroxyvitamin D2 and 24,25,26-Trihydroxyvitamin D2: Novel Metabolites of Vitamin D2+

G. Satyanarayana Reddy\* and Kou-Yi Trerngs

Department of Pediatrics, Case Western Reserve University at Cieveland Metropolitar General Hospital, Cleveland, Onio 44104, Department of Pediatrics Brown University Program in Medicine, Women and Infants' Hospital of Rh. de Island, Providence, Rhode Island 02905, and Department of Pharmacology. Case Western Reserve University and Veterans Administration Medical Center, Creveland, Onic 44106

Received Jul. 19, 1987 Fevised Manuscript Reseived August 29, 1989

ABSTRACT: Understanding of the inactivation pathways of 25-hydroxyvitamin De and 24-hydroxyvitamin D<sub>2</sub>, the two physiologically significant monohydroxylated metabolites of vitamin D<sub>2</sub>, is of importance, especially during hypervitaminosis D<sub>2</sub>. It a recent study, it has been demonstrated that the inactivation of 24hydroxyvitamin D<sub>2</sub> occurs through its convenien into 24.29-diaydroxyvitamin D<sub>3</sub>. Thoszewski, N. J., Reinhardt, T. A., Napoli, J. L., Beitz, C. D., & Horst, F. L. (1988) Biochemistry 27, 5785. At present, little information is available regarding the inactivation pathway of 25-hydroxyvitamin D<sub>2</sub> except its further metabolism into 24 28-dinydroxyvitamin D. Genes, G. Ecsenthal, A., Segev, D., Mazur, Y., Froicw, F., Halfon, Y., Rapinovich D., & Shakked, Z. (1979) Bischemistry 18, 1094]. In our present study we investigated the metabolic fate of 25-hydroxyvitamin  $\Gamma_2$  in the isolated perfused rat kidney and demonstrated its conversion not only into 24.25-dihydroxyvitamir. De but also into two other new metabolites namely 24,25.28-trihydroxyvitamin D<sub>2</sub> and 24.25-26-trihydroxyvitamin D<sub>2</sub>. The structure identification of the new metabolite. was established by the techniques of ultray clet absorption spectrophotometry and mass spectrometry and by the characteristic nature of each new metabo, te's susceptibility to sodium metaperiodate oxidation. In order to demonstrate the physiological significance of the two new triby irony metabolites of vitamin Dz. we induced hypervitaminosis  $D_2$  in a ratius ng  $[1\alpha^{-3}H]v$  tamin  $D_1$  and analytical its plasma for the various [30-17] vitamin De metabolites on two different high-pressure liquid phromutography systems. The results indicate that both 24,25,28-trinydroxyx terrain Dound 24,25,26-trinydroxyxitemin Docirculate in the vitamin De intexticated rat in sign ficant amounts along with other previously identified monchy froxy and citydroxy metabolites of vitamin D<sub>2</sub>, namely 24-h drowy itamin D<sub>2</sub>, 25-hycroxyvitamin D<sub>3</sub>, and 24,25-dih; drowy itamin  $D_2$ . Thus, it may be hypothesized that the two new trihydroxy metabolites of vitamin  $D_1$  play an important physiological role in the ceactivation of 15-hydroxyvitamin Dy, especially during hypervitaminosi; Dy

Chiracter Vitamin Do (ergoculciferci) and vitamin E (thelesuloriero) with the fare the two important natritional form of vitamin E out of the lesuloriero in the land of the l  $oldsymbol{V}$ itamin D $_0$  (ergoculciferci) and vitamin E  $_0$  (chelebulorfaro ). suda est the known forms of vitamin D. Napoli et al. 1979.

Suda est Aprovitamin E. 7-dehydrocholesterol) exists in the kin of many ower organisms such as fungiant years. Frontamins D. and E. are converted into their orresponding vitamin. primite Dyand by when exposed to UV radiation. The two vitamins and differently in the structure of their side chains (the lide chain when the books of extra method eround to C-24 and a double for Giffer only in the structure of their side chains the lide chain  $E_0$  vitamin  $D_0$  has an extra methyl group at C-24 and a double bond between C-20 and C-23 when compared to the hose chain geof vitamin Di) Historically, vitamin De became important atin clinica med cine as it was the first synthetic v tamin D preparation available for the treatment of richets, and this still being widely used to satisfy both therapeut it and natritional preeds of man and other commercially important man mals.

. S. ir

At present, it is the general belief that the further metabolic pathways of vitam n Do are similar it those o vitamin Do (Norman et al., 1982). Vitamin D., ille vitamin I., undergoes hydroxy attonicat C-23 in liver and at C-1 in kidney to form 1.25(OH), D2: the Fermonally active form of viramin D2 (Jones et al., 975). During the past decade, the pathways of side-chain metabolism of vitamir  $D_2$  n etabolite (13-OH-D) and [25(OH, 2D]] have been studied extensively. It is now appearent that the lice chains of both 15-0M-Dy and 4.25-(OH 5D) undergo and ogous metabolic alterations resulting in the formation of many relatively mactive metabolities, and this subject has been extensively studied in several laboratories and was reviewed recently by Jones et al. (1987). Because of the obvious structural difference; be ween the side chains

Appresiations 24-OH-Ds 24 ayers with min 2: 25-OH Ds 25-

A borevistoris i Le-Omino, Le Rydrox vivismin 15, 17-Omilog, 11-7 (droxyvitamin 15, 14-25) OHI<sub>2</sub>D<sub>1</sub>, 24-27), 10-diaperc vivinamin 15, 10-20-OH <sub>2</sub>D<sub>1</sub>, 24-25-34, 25-34,

24,25, Tetrion drogsystation De 1,25,0Hi,De 1,25-drogst Kipytamin De

USDOWN THIRD GROWN LATTING OF THE CONTROL OF THE CO

UNA 1-246 - Project 40-423 (15 f)

e andres simulates acres our maineauth EXHIBIT B

Partions of this work were presented at the Principle for the Conubling Lawrence of this work were presented at the Profindernation. Con-tropic ference on Calcium Regulatine Hormones and Bone Metabolism. 950. Abuse. Fronte, and at the Hurrial Meeting of the Suciety for Profiatric Lawrence, 1957. Anulism, CA. Tolk work was surported to burn by a Brant DK-30128. From the National Institutes of Health to D.S.P. Une Women and Interns Hospital of Royde Isla d. 177 I usie, St 森 Mis Wigner and Iola 第: Picylainee, R.1 0297.1

Gase Western Resente Unit ersity at Tieveland, withoppisit in Fierera fig. 1988 Western historic of Layer and St. Royde Island Again para and Wamen and Infanta Hospita. 31 Royde Island The Commission of the C

<sup>1</sup> Case Western Heserie University and Venerals Administration

of vitamins  $D_{\xi}$  and D , it appeared logical to find different pathways of a de-chair metabo sm for vitamin De metabolites  $[25\text{-}O\hat{H}\text{-}D_2]$  and  $1.25(O\hat{H})_2\hat{D}_2$  . Accordingly, a novel metabolic path va for 1.25 OH), D; has been recently iescribed, and it has become opyous that the side-chair metabolism of  $1.23(\mathrm{OH}_{\odot}\mathrm{D}_{2})$  differs from that of  $1.25(\mathrm{OH})_{2}\mathrm{D}_{3}$  and that the differences become apparent on a after the C-24 hydroxylation step (Herstler al., 1986, Reddy & Teerng, 1986), Fishowing The C-24 hydroxylation step. .25(OH 2D; undergoe: hyprobabilitions either at (2.28 to form 1.24.25.28 OH 1.De or at 13-26 to Nom : 14,25 16(OH), Do (Feddy & Tserry 1986). Even though a complete functional evaluation of the two new metab. Ref. of  $-.25({\rm OH})_2\mathbb{O}_2$  hat not been performed yet, it appears from our presentery result that the new puthway coside-thair metaboli miof (,25) OH (<sub>2</sub>D<sub>2</sub> is indeed a fathway intended to inactivate (  $25(OH)_{\rm T} {\rm D}_{\rm S}(G,S)$  Relay on t  $R,\, \Sigma_{\rm S}$ Horat, aroub shed opervations). At present, very little is known regurding the inactivation pathway of 25-0H-III except its ochver yor (to 24(P) 35(OH); Ut (Jones et al., 1974) 1980). We predicted the inactivation of IT-OH-Derbrouch its conversion into 24.25.28 OH: D: and 04.25.25.(OH) T: in a manner and ogcus to the mactivation of 1.25(OH 3F; through it) conversion into 1.24.25.26 (OH  $_2D_2$  and 1.24.25.26 (OH)  $_4D_2$ Therefore, this study is performed with the nim to scentify 24.25.29(09H),  $D_{\rm b}$  and 14.25.26  $\rm (OH), D_{\rm b}$  as the further mass tabolite, of IS-OH-D; and to demonstrate their formation er pecialty during hypervitamine is Di-

# MATERIALI AND METHODS

General. U traviolet absorbands spectra were tiver in 2-probation with a Beckman DNL a recording spectraging continue eter High-performance liquis entima ography (HFLV) was performed with a Waters Model 6/30 equipped with a perector (Mode 440) to monitor UV-absorbing material at 174 am (Waters As bounter, Mirford, MA - ALTHPLC relivery, were purchased from Burdick & Jacks, a Laboratories Michega. MT Miss spectra (70 eV) were obtained on a Heislett-Popkard 59-5 B mass absolutionness. Samp es of me abolities (0.5 agleser) were introduced into the for source maintained at 200° PD vala direct-insertion probe

Pitam n.D. Tempounas - Vitam n.D. was purchaled from Sigma Chemical Co. (St. Leads MO), 25-0H-Fg was a gift from Drs J. A. Campbell and J. Baboock (Upronn C. . Kaiamazoc,  $M(r) = 24(R).25(OH)_{\rm C} T_{\rm c}$  was a gift from Eq. T. Konsyashi, Floor Women's College of Pharmacy, Kobe, Japan. All the various symbolic standards of both vitaming Es, and D<sub>3</sub> metabastes used in this study were a gift from Ers. Milan Uskokovie and E. C. Baggiolini (Hotfmann-La Roche Inc., Nurley, NJ) Authentic 24-OH-D; was isolated from the serum of ottam r. Do intoxicated rate, and the structure of biologically chodined 24-0H-D  $_{\rm 2}$  kis varified by the a specified trometry us described before (Jodes et al. 1985). [2009H]-Vitumin De (1.2 Clemmo: was a gift from Dr. J. L. Napoli Guste University of New York, Buffale, NY

Antimais. Male Sprague-Dawle (about 310 g. purchange from Zivic Miller Laboratories, Inc., Allison Park, PA. werd let a regular rodent blet (Alg way, byracuse, NY), containing 5,977 cast am. 3 85% phosphorus, and vitumin D. 1045

Study of 15-014-0; Merabilism in the hidney of Filantin De Intoxicated Rais Laing he Technique of Klanes Perfusion. hildney perfusions were performed as described perfore in detail Reddy et al., 14-2, 1963 - In order to produce the various arthur metacolities of 254 (H-Dy In submittees sufficient for ne r structurel (contification, we performed two kildnes berutilist il has been hombo possible been.

1,954) that the activity ( the enzymes involved in the fundamental of COL 1, and 24.25 OUT. metabolism of 21-OH-L and 24.25(OH). De can be indicated the control of the contr in the k dneys by intoxicating the rate with vitamining the Therefore, in these perius on experiment, we induced to pervitam notif D- in rate to order to increase the and pervitam notif D. in rat, in order to increase the activity the the renal enzyme, that are involved in the further metability. of 25-OH-D. On the ball of the information given into previous itudy of Shepard and DeLuga . 480), we first timated 1000 ug of vitamin D<sub>2</sub> at a safe total dose that the intoxicate a rat without causing death. We then induce hypervitaminosis D<sub>2</sub> in each rat by adminitering 1(n) ug vitamin I - in 100 all of 95% ethanol ir tramusculariv acti day for a period of 10 days. The last dote of vitamin D. administered to the rat 24 p propriet the solation of the edge. Each kidney a stated from a sitamin De intexicated rat with perfused for 4 n with cold 25 OH-Pt 200 nmoi in 100 m of ethanol) which was introduced into 100 mL of perfusally after a 5-mir abilization period following the isolation the kidney. In this study, we also performed a control ? fusion experiment in the absence of a kidney and demonstrated that there was no metabolism of 25-OH-Ly in the perfusion Eppuratus in the absence of a kidney (fare not shown in the appearance of the not shown in the appearance of the sidney partial processors.)

Lord Estraction. Lip c extraction of the ordney perfusation of the was performed upported by the procedure of Bilgh and Dyer was performed upported by the performed was substituted to first eluter chloroform.

25-CH-D: from Kidney Perfacate for I me. 22.

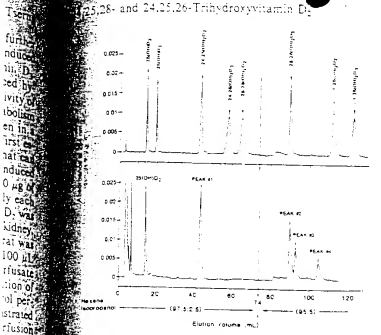
26-CH-D: from Kidney P Isolation and Furification of the Various Metabolices of the cane 25-CH-D. From Kidney Perfacate for Their Structure idea in more point of the Angle extruct absained from \$60 mL of the Chichero jected directly to HPLC under the chromotogruph a condition described in the legend to Figure 1 without ever bading we occlumin with about. Even though radiotable'ed 25-OH-D, without available this satisfie time of our present study, we went about a trace the larious metabolites of 15-OH-D, by months. described in the egend to Figure 1 without ever pading its toring their UV upsorbance as 204 nm. The e.u. on volume of such UV pear curing the first HFLD system shown Figure 1 (lewer pune ) was a (slidws peak 1 42-46 mL; peak 1. 88-92 mL: neuk 2. 92-96 mL: peak 4 (44-)63 mL. Athe sciented UV neaks obtained during the limit rune HPL rans were too as and each individual UV reak was the subjected to a scotted HPLC system with the same Zorbax-SI column (4.5 mm x 25 cm) eluted with metry endichloride collated f 2-programol (94 6) at a flow rate of 1 mL min. The elutio condition volume of each C+ peak during the second HPLC system with a funding a formula peak. ar follows | peak | 12-14 mL; peak 2 30-3; mL; peak 24-28 mL, peak 4 4)-48 mL. Each UV yeak ibtzined from the first HPLC run. At this time, the purity of each UV per was tested by obtaining their UV spectra. The UV shorbing the majority of run per up the majority from page. was tested by obtaining their UV spectra. The UV-absorbin 24.25(C) material from pears - 2 and 4 exhibited a UV-absorbing metabatic is character stic for all D vitamins. The UV-absorbing material to a spectral description of the UV-absorbing material to a spectral description. terial from peak 3 exhibited. L.V. protrain with a UV and 243 nm and thus did not exhibit a L.V. spectrum characterist. too D vitam ns. All inree vitamin D<sub>2</sub> metabolites obtains. for D vitam ns. All incree vitamin De melabolites obtains from beaks. It and 4 were then subjected to odium metabolites also periodate by dation and mass spectrumetry in order to identify as also their chemical structures.

Sodrum Metapermoute Octidation of 24,25, 019,5- and the The New Metapointer of II-DE-D. The susceptibility 24.25) OFF 2D2 and the two new metapolities of 15-OH-Di Aldrum metubeniboate (Nat Callow at 1.2 allow was tosted in one to south the exact positions of the hundry of orongs in each

Carried c 解HPLC () otall thr 0 Figure ponding (OH)<sub>3</sub>D incubation difference between

ninderec dation. Stuay

af Cas



ed:

ivity of

ibolism

en in

1751 6 nat cal

nduced i

0 μg of

D. Lidney

di Was 100 山東

riusate ...

ol per

[ . was ..

ax-SILin.

Bullion .

TT #35 \*\*

rear 3.3

i from ]

to with

vineak,

orting

m .ha!

ig Fier

, <sub>ma</sub>, it

teristic

tained

~e.2

jenu y

... Isf

.gi<sub>=</sub> : a

order.

ೀಚಿ ಪ್ರ

فاررد

or de

riusion (a) Signature of a maxime of various authentic synthetic riusate (b) Dyer (b) The cares of Dyer (c) The care is performed on a Zorbak-Sil column (25 cm × 4.6 mm) that was office eluted with hexane-2-propanol (97 5.2.5) at a flow rate of 2 Limin until 25.28(OH), D. was eluted out of the column. Then, the first solvent system was changed to a second solver; system hexane—2-propanol. 95:5), beeping the flow rate same to elute the elder the increase metabolites of vitamin Doot of the column. The various of the metabolites of 25-OH-D, were identified by monitoring their LV as was a labor band and the column. Fears 1.2 and 4 represent 24.25(OH), Is, and 24.25.25(OH), Is, and 24.25.26(OH), Is, ns was 24.25 28(OH)<sub>1</sub>E<sub>2</sub>, and 24.25.26(OH)<sub>3</sub>E<sub>3</sub>, respectively. Peak 3 represents a non-vitamin D contaminant produced by the kidney. Note official distance of the 25-0H-D<sub>2</sub> peak shown in the lower pane of the ing the figure represents only  $\gamma_{10}$  that of the wriginal peak. D<sub>2</sub> was  $\gamma_{10}$  was

Do was a between two carbons when either both carbons bear hydroxyl monitor groups or one carbon bears a hydroxyl group and the other solume? I bears a keto group. Each metabolite (0.3-0.5 µg each) was own in dissolved in 15 µL of methanoi and was allowed to react with peaking 10 µL of 5% aqueous NaIC4 for 5 min. All the reactions were L. Allow carried out at room temperature (25 °C). The appropriate HPLOs systems used to isolute the periodate cleavage products sinen allows a librar metabolites are described in detail in the legend of all three metabolites are described in detail in the legend to Figure 2. Even though all three metabolites of 25-OH-D. isolated from the kidney perfurate were susceptible to periodate oxidation, we noticed that during an incubation period of 5 min only 5% of 24.25 (CH D), was converted into its corresponding periodate cleavage product. whereas both 24.25,21- $(OH)_{i}D_{i}$  and  $14.25.26(OH)_{i}D_{i}$  were completely converted into their corresponding periodate cleavage products (Figure (£2). In order to produce the periodate cleavage product of

24.23 OH De in sufficient quantity, we had to increase the incubation period to ! he duta not she was. The reason for the differences in the degree of asceptibility to periodate ox cation between the three metabolites of 25-CH-D<sub>2</sub> s as follows. The twicinal diol at C-24 and C-25 in 24,25+CH<sub>2</sub>D<sub>2</sub> is not readly suscentible to periodate exidation and it is sterically hindered This C-18, C-27, and C-25 methyl groups. This phenomenon alwas a to noticed by Jones et al. (1979). However, the Acinal Goldina C-24 and C-25 in 14.25.25 OH (P) and the vicinal diological C-25 and C-26 in 24.15.25 OH (P) are not stee cattle C-25 and C-26 in 24.15.25 OH (P) are not stee cattle C-25. Adindered and, hence, are meanly susceptible to periodate twodation

hastade general active abouter of country De this Vistamin The Control of Area Rule Was experiment was designed by semonstraine the in vivil extenses of two new triplycously metalwings.

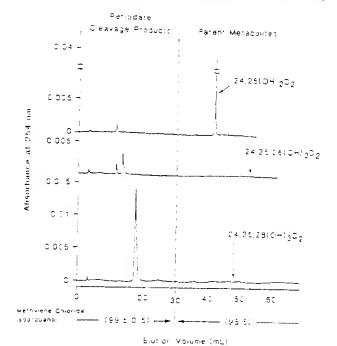


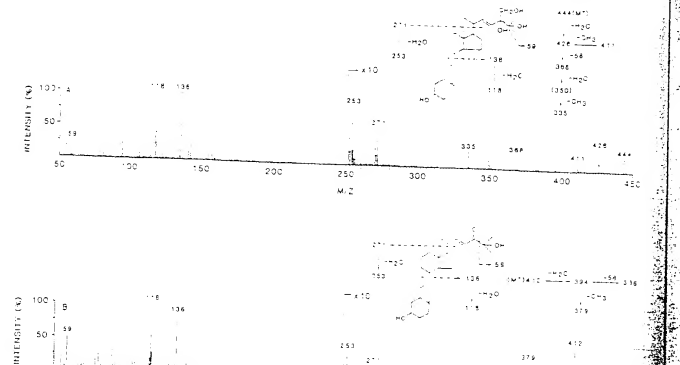
FIGURE 2: HPLC analysis of the reaction products, obtained by treating 0.3-0.5 µg of each metabolite of 25-OH-D, with sodium metaperiocate for 5 min 24,25(OH)<sub>2</sub>D<sub>2</sub> (upper panel); 24,25,26-(OH)<sub>2</sub>D<sub>2</sub> (middle panel); 24,23,23(OH)<sub>2</sub>D<sub>2</sub> (lower panel) HPLC was performed on a Zorbak-S.I column (25 cm < 4.6 mm). The column was first eluted with methylene chloride-I-propanol (99.5:0.5) at a flow rate of 2 mL/min until the periodate cleavage product(s) of each metabolite eluted out of the column. Then, the solvent system was switched to methylene chloride-2-propano. (91.5) at the same Now rate to eluce the unreasted parent metabolites. Arrows incleate the elution position of the parent metabolites.

of sitamin D<sub>2</sub> in hypervitaminesis D<sub>2</sub>. Because of a limited supply of 3a-3H vitamin D3, we only performed this experiment in a single rat. Hypervitaminosis D; wa induced in the rat with the same dose of vitumin D2 and the method that has been used in our present study during the investigation of in vitro metabolism of 25-OH-D2 in kidneys isolated from vitamin  $\mathcal{D}_{z}$  intoxicated rats. We first prepared a mixture of nonradisactive vitamin  $D_1$  ( 000 µg) and  $[3\alpha^{-3}H]$  vitamin  $D_2$  (20 gC) in 1 mL of 95% ethanol and thus obtained a specific activity of 20 cpm/1 ng of vitamin Dis. The rat received 100 AL of the above mixture intramuscularly each day over a per od of 10 days. Twenty four hours following the final cose, the rat was sacrificed by excanguination to obtain 6 ml of pia ma, which was divided into two portions. The first 3 mL persion was again divided into three .-mL portions. All four plasma samples were extracted, and the lipid entract of each sample was subjected to HPLC directly. We first performed pre iminary HPLC runs of the lipid obtained from 1-mL plasma samples using the HPLC system described in Figure On the basis of the information obtained from the three preliminary HPLO runs, we developed the HPLO system described in the legend to Figure 5 and measured the various [fir]vitamin D<sub>2</sub> metabolites present in the final 3-mL piusma sample.

# RESULTS

Metabolism of IS-OH-Eq by the Perfused Ridney sociated trom Vitamin  $\hat{\mathcal{O}}_{\mathbb{C}}$  intoxicated Ratio. The light contentiate obtained from 20 mil of perfusate was analyzed in a single HPLC run on a straight phase HPLC system (Figure - upper punell that a canable of resolving most of the known mewoodten in beth 25-0H-C land 25-0H-C, From the HPLO

50



253

250

M/Z

FIGURE 3: Mass spectra of 24,25,28(OH)-D2 (A) and it, periodate cleavage product (B).

200

150

chromatogram in Figure ! (lower pane ), it became apparent that there were only four UV peaks (peaks 1-4) following the UV peak of 25-OH-D2. Out of the four UV peaks, UV peak I was found to be a non-vitamin D lipid contaminant. UV peak 1 comigrated with the synthetic standard of 24(R), 25-(OH) Do on two different HPUC systems and exhibited a mass spectrum identical with the one described previously by Jones et al. (1979-1930) (cata not shown). Also, the metabolite was susceptible to periodate exidation and resulted in the formation of 24-Neto-25,26,27-trinor-D<sub>2</sub>, the expected periodate cleavage product of 24,25 OH 2D2. The mass spectrum of 14-ketc-25,26,27-trinor-D; was identical with the one previcusiy described by Jones et al. (1979) (data not snown). Thus, on the basis of the periodate oxidation and the mass spectrametric data, the metabolite of 25-OH-D $_2$  in UV peak I was identified as 24,25(OH)2D3. The metabolite of 25-OH-D3 in UV peak 2 comigrated with the synthetic standard of 25.25-(CH), D; (Figure 1) and was later identified an 24 25,23. (CH), D. The metabolite of 25-OH-D2 in UV neal 4 migrated just before the synthetic standard of 1.25(CH)<sub>2</sub>D<sub>2</sub> (Figure 1) and was later identified as 24.25.26(OH), Dp. Thus. the results of our study indicated that 25-OH-Dig wits metabolized in the isolated perfused rat kidney into three major metabolites of vitamin  $\hat{D}_2$  but of which  $24.25(OH)_2\hat{D}_2$  was described originally by Jones et al. (1979, 1980) and the remaining two metabolites were found to be new. The detailed description of the structural identification of the two new merapolities of 2340H-D<sub>2</sub> as 24,23,28 OH <sub>3</sub>D<sub>2</sub> and 24,23,25-OH De is as follows

100

Structural Taent fination of 24,25,2800H D- and 24.25.256.044  $_3\mathcal{D}_2$  . The  $_{1,k}$  -new imeraposities of puraled from the kiches herbisate, exhibited UN spectra with un absorbance mus, mum a Disse nom and an absorbance minmum at 228 am. This "calog indicated that the twic metabelities contained un initiam con-literimene chromophere i déta

not shown). The may spectra · Figurer 3A and 4A) of the new metabolites eith titled peak, at m/z 271, 252, 136, and 113. Collectively the peaks indicated that the secosteroids nucleus of their parent 25-OH-D; has remained unchanged and that the two new metapolites were formed as a result of changes occurring on vion their side chains. The molecular ion at M = 444 (M) i.: the mast spectrum of each new me tapolite indicated that noth new metabolites contained two additional hydroxyl groups when ecopared to 15-OH-D<sub>2</sub>. As we had thready determined that the secosteroid nucleus of both new metapolities was indict and similar to the one present in 25-OF-D<sub>2</sub>, it was pishole to conclude that each new metable cite was formed as a result of audition of two hydroxyl groups to the rive chain of 1500H-D2. The exact locations of the two additional hydroxy groups on the side chain of each individual new metabolite were determined in the following way.

336

350

400

300

The mass spectrum F gure 3A) of the new metabolite in UN peux 2 in Figure 1 exhibited a beak at myz 59 and a peak at m = 356 formed a. . result of elimination of 58 mass unit from the peak at m is 41. (Mollafferty type rearrangement) This fir ding indicates that the met spoilte contained an inter-C-25 hydroxy group with no hydroxylations recurring on C-26 and C-27. Also, this metabolite was susceptible to periodate oxidation, and the mass pectrum of the periodate cleava. product i mgure 3B) shi wod a molecular for at m = 412 which indicated that the parery compound had lost CH3OH (32 mag) units during the process of persocase exidation. The charge actenistic peak at m. r. r. and a peak at m. r. 336 formed a result of alimination of . 8 mass unit. from the peak at me - Mallafferty tyre rearrangement indicated that the periodate sleavige product still contained an intact C-25. utilin the remodule orealinge product was identified 5.82784 - 14-14811-13-15 (1-12) Nie man observed that 25.0U 24-x std-28-0 ardDe was further baseentible to periodate 🕬

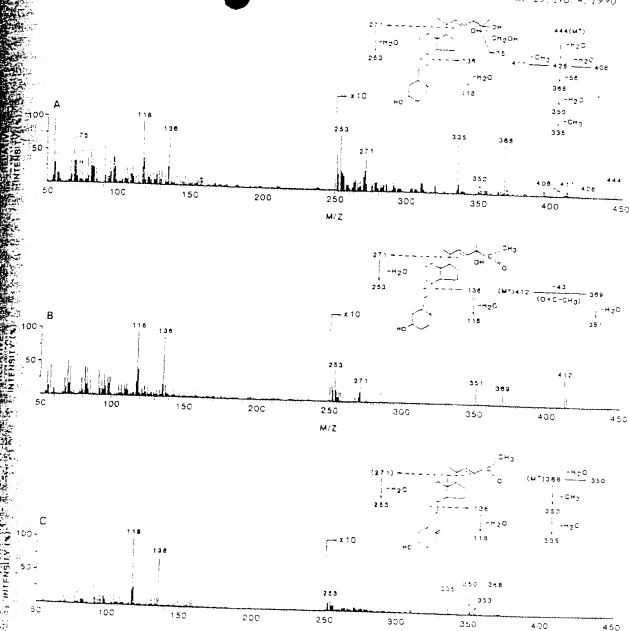
100

24,25,25-

INTENSITY (%)

INTERSITY (%) RELATIVE

Cation, a OH-24. that he With an e



**COURT 4** Mass spectra of 24.25.26(OH)<sub>2</sub>D<sub>2</sub> (A) and its two periodate cleavage products ( $b \cos 2$ )

edition, and this finding provided indirect evidence for the retion, and this innuing provided indicated and hydroxyl described at C-251 in 25-OH-24-keto-28-nor-D<sub>2</sub>. However, we must at C-251 in 25-OH-24-keto-28-nor-D<sub>2</sub>. However, we mit soliate the C-24 acid, the expected periodate cleavage in D-25 as the HPLC systems eccloped in our laboratory at the time of this study were not Place to iso are highly polar acids. The formation of 25-OH:24-18to-28-nor-Do from the new tribydraxy metabolite Sylvamin D. as a result of periodate exidation would be possible only of the new metabolite contained vicinal hydroxyl pupps a C-24 and C-28. Thus, it was finally concluded that take the new tribyerbay metabolite of vitamin De possessed by-Toxyl groups at C-24 and C-28 in addition to the original by Ydroxy group at C-25, present in its parent 25-0H.D., and Yas the office identified as 24.25.25(CH.D.)

26 late

25

- 11

The mass spectrum (Figure 4A) of the metaholite in UV Active in the specifical to again a large of the staggested halt the inetage are contained an anactive of the specifical format (2-25 by 350x). Showing the specifical format (2-25 by 350x). Hat the metable sie contained an illiad. O I in order convincing the an extra of driving group at C-26. Futner more convincing to the more convincing of the Fridence of Chin East No. la lugi dame que to in e presence of

characteristic mass flaements at  $m_{e}z$  363, 350, and 335. These mass fragment - vere produced as a result of McLafferty type rearrangement ian  $\alpha$ -substituted  $\beta$ -hydroxy aldehyde resulting from the deny fration of the molecular ion  $(M^{+})$ . followed by a loss of OH3CH3CHO (58 mass units). This phenomenon is similar to the characteristic decomposition pathway described for prisupportuned 3-hydroxy esters (Budzikiewicz et al., 1967 - First termore, this new metabolite was also susceptible to pur ocate oxidation and gave rise to two cleavage products (Figure 2). The mass spectrum of the more poier periodate cleavise product (Figure 4B) showed a molecular ion at m z 4/2 wh or indicated that the new metabolite had lost CH<sub>2</sub>OH -31 maps units aduring the process of perlocate objection. Also the churacteristic loss of  $CH_1 + C = 0$ (43 mass units) from the molecular ion gave rise to the peak at m = 369 . Which this information, the more polar periodate ofearrage or waser was licentified as 24-OH-25-kete-25-nor-Dethe formation of which while only be result to the nnesence if moine, in it my groups at 0-25 and 0-26. The ess pertir her, some e ear age product had exhibited scentical

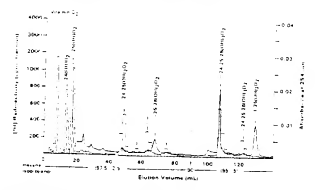


FIGURE 5. HPLC profile of the lipid extract of a plasma sample 3 mL+ obtained from a rat given 1000  $\mu g$  of [3H] vitamin  $D_{\tau}$  in divided doses over a period of 10 days. The plasma sample was mixed with authentic nonradioactive v tamin D<sub>2</sub> metabolites (24-OH-d<sub>2</sub>, 0.2  $\mu$ g; 25-OH-D<sub>2</sub>, 1.2  $\mu$ g; 25,28(OH)<sub>2</sub>D<sub>2</sub>, 0.2  $\mu$ g; 24,25,28(OH)<sub>3</sub>D<sub>2</sub>, 0.5  $\mu$ g; 1,25 OH <sub>2</sub>D<sub>2</sub>, 0.2  $\mu$ g) with the aim of identifying each individual radioactive viramin Do metabolite by its comigration with its corresponding authentic nonrad oactive vitamin Demetabolite standari. The lipic extract of the plasma sample containing the various [3H] Atamin D<sub>2</sub> metabolites and the authentic nonradicactive vitamin Di metapolite, was analyzed by HPLC under the chromatographic conditions described in the legend to Figure I except that the first so rent system was changed to the second solvent system at an elution volume of 90 mL. The elution positions of the various horradioactive authentic standard of vitamin D<sub>2</sub> metabolites as monitored by their UV absorbance at 254 nm are depicted in the figure above by the solid line. Note that the UV absorbance profile during the first I0-min HPLC run is not shown for the sake of clarity. Fractions of mL during the first 10 min of the HPLC run and fractions of 2 mL during the remaining per oct of the HPLC run were collected. The HPLC efficient in each fraction was divided into two equal portions. The first cortion was used to measure the radioactivity, which is depicted in the figure by a histogram. The second portion was used for recommandgraphy of each major metabolite of vitamin Lip on a second HPL/C run using a methylene co.oride-2-propanol mixture as the solvent system (data not shown). Thus, we confirmed the identity of each major metabolite c. vitamin D. by its comignation with its corre bonding authentic cold standard on two different HFLC systems.

chromategraphic mobility as that of the periodate cleavage product if 24,21:OH). D2 (Figure 2). Its mais spectrum (Figure 4C) was identical with the mass spectrum of the per odate cleavage product of 24.25(OH-2D2, previously published by Jones et al. (1979). With this information, the less polar cleavage product was identified as 24-keto-25.26.27-trinor-E(2). The formation of 24-keto-25.16.27-tri $ncr \cdot D_2$  from the new trihydroxy metabolite of vitamin  $D_2$  as a result of periodate oxidation would only be possible if the new metabolite contained vicinal hydroxyl groups at C-24 and C-25. Thus, by putting together the aforementioned data, it was finally concluded that the new trinydroxy metabolite of vitamin D<sub>2</sub> possested hydroxyl groups at C-24 and C-26 in addition to the original hydroxy, group at C-25, present in its parant 25-OH-D<sub>1</sub>, and was therefore identified as 24,25,26-

luentification of both 24,25,28(OH), D. and 24,25,26-Girando as the in Vivo Metabolites in a Vitamin L. Intoxscated Rat. From the HPEC chromatogram shown in Figure 5. it became obvious that there were several orbulating (Figure men D) metabolites in the prasma of a vitamin  $D_{\sigma}$ intoxicated rat. We were able to estimate the concentration of each withmin Dometabolite as we knew the specific activity of  $[^3H]_{\rm Violamin}$   $D_2$  that was administered to the rati. The mean value of each metabolite concentration in 1 mL of plasma estimated from two different HPLC runs was at follower mumin D<sub>1</sub>, 438 ng: 24-0 H-D<sub>2</sub>, 39 ng: 23-0 H-D<sub>2</sub>, 96 ng: 24.25 OH 305, 28 ng: 24.25.25 OH 305, 31 ng: 24.25.24 OH 305, 13 ng Thus, from the results of our sarefully performed HPLC analysis of the clasmo of the vitamin De

interviented rat, we established so th 24.25.28 OH 13D, and 24.21.25(OH), D<sub>2</sub> as the agricultant in vivo metabolite, in the special standard of the pervitaminosis D<sub>2</sub>. Also, our firding of both 24 OH D<sub>2</sub> and before of pervitaminosis D<sub>3</sub> was not curprising in light of a recent study. Four preserving the standard of the pervitaminosis D<sub>3</sub> was not curprising in light of a recent study.

hypervitam most D<sub>2</sub>. Also, our firding of both 24-OH-D<sub>2</sub> and 24.27(OH<sub>2</sub>D<sub>2</sub> as the major circulating meta-clites in hypervitaminosis D<sub>2</sub> was not surprising in light of a recent study by Kostew ki et al. 1983. We also noted in Figure 5 that there were one major and two minor radiouctive vitamin D<sub>2</sub> in 25.26(O) meta-order on the minor adiouctive vitamin D<sub>3</sub> in 675.26(O) meta-order on the intention of the meta-order peaks, it could be predicted that one of them especially the major one might be 24.25(OH<sub>2</sub>)<sub>1</sub>, a neta-order that was recently identified by Kostew ki et al. 1.983 at one of the major circulating meta-order of vitamin D<sub>2</sub> in radiomexicated with vitamin D<sub>3</sub> Considered.

This paper reports the itentification of two new meta-boiling of 31-O(H<sub>2</sub>)<sub>2</sub> produced in a mamoralian kidney. They were donarised at 24.25(2)(OH<sub>2</sub>)<sub>3</sub> and 24.25(2)(OH<sub>2</sub>)<sub>4</sub> in the vitamin D<sub>3</sub>. The produced in a mamoralian kidney. They were donarised at 24.25(2)(OH<sub>2</sub>)<sub>4</sub>, and 24.25(2)(OH<sub>2</sub>)<sub>4</sub>D<sub>3</sub> and Co<sub>4</sub>Co<sub>4</sub> to 10 of 21 OH<sub>2</sub>C<sub>2</sub> was identified at Alace at the one we meta-boiling of 21 OH<sub>2</sub>C<sub>3</sub> was identified at Alace at the one we meta-boiling of 21 OH<sub>2</sub>C<sub>3</sub> was identified at Alace at the one we meta-boiling of 21 OH<sub>2</sub>C<sub>3</sub> was identified at Co<sub>4</sub>C<sub>4</sub>C<sub>5</sub>C<sub>6</sub>OH<sub>2</sub>D<sub>3</sub> and a 24.25(D<sub>6</sub>OH<sub>2</sub>D<sub>3</sub>) in this study, we only demonstrated the formation of the rew in eta-boiling at administration of the country of the country of the country of the rew in eta-boiling at administration of the country of the rew in eta-boiling at administration of the country of the rative study that followed our arcoent study, we first synd the inedicine a theorem out H + 25-C/H+E<sub>2</sub> versy mutically by perfusing livers to the inedicine a thesized out the LD-Circ-Lip way manusching open a band manusching in a 225-OH-D. Located in the vitam in D cell a entire tis with [33--H] vitaming a 225-OH-D. Lip. We then demonstrated both 24.25.2-(OH)<sub>3</sub>D<sub>2</sub> and a collection in 24.25.15(OH)<sub>3</sub>D<sub>4</sub> as the phy wiegonal metabolities of 25-OH<sub>2</sub> and the high 24.15.16 (OH) 30; as the phy allogant metaborites of 15-OH; to the high by versus agriculture train normal rais on a regular fig. 1985). In a codent diet with a physic agreat a noentration of 13a-1H] the metaborites of the physic agreat a noentration of 13a-1H] the metaborites of the physic agreement of the previous of the physics of are unable to assert the bicker of outility of outil 24.25.28 D intoxica (Ciris Dy and 24.15 21 (OH) Dy and to the unavailability of the converse these new metabolites in a quantity sufficient for the standard b on save mentaring intestinal color are transport and boness. A pathway of colorair mobilization. However, calcium motifization. However 2. in alternative, tis still to the Jones et a pons be for us to present that bein 24.25.25(OH),D. and the lactivation 21-CH D. as we have recently bound that the two leads of Thus, the hydrox-lated retabolites of 22 CH D. as we have recently bound that the two leads of Thus, the hydrox-lated retabolites of 22 CH OH), D. and 24,25,26 Church (OH), D. [1.14 25 CS (OH), D. and 24,25,26 Church indeed completely nature. indeed completely inactive to learn or intertransport and bone calcium mobilization (G. 5) Fleddy any
R. L. horst, incuratehed of servation). Furthermore in out.

present study we have demonstrated that both 24.25.28 the difference of the complete of the difference of the complete of (OH) Do 11.14 25 28(OH) Do and 24.27,26(OH) Do and indeed completely inactive to error of intertinal calciums two rew tribydraky metabolites of atomin by it appear logical it the present time to us ame that the formation of never metabilites may play an important physiological role in the deact vation a 25-Gh-D; express, a saring hyperv taminos

tabolism of 25-0H-By true 25.28 CH <sub>2</sub>D<sub>1</sub> and 25.26-OH)<sub>2</sub>D<sub>2</sub> We first demonstrated that there was no metade them of 25 OH-Do no 1 .28 OH (Do no the twitted herfuled kidney) as indicated by the absence of a fix-absencing peak in the migration position of the authentic lynthatic tandard of a 25.25. CH 17: Figure . . La er. we also demonstrated that it

oiles of 25

Blochemy

nin **D**. ibolile y west bolin in our

ney 5 the

-d. d.

min D the att it court unites mili

ing metabolite of 25-OH-D<sub>2</sub> in rat. Thus, it appears that preferred substrate for the entry med responsible for both 28 and C-26 hydroxylations is 24,25, OH), D2, but not 25u:auni sHD2 In our previous study (Redcy & Tserng, 1981) we The design of the state of the decay at Iserng, 198t) we we demonstrated that 1.25(OH)<sub>2</sub>D<sub>2</sub> is hydroxylated first at 24, to form 1.24,25(OH)<sub>2</sub>D<sub>2</sub>, which is then further hydroxylated either at C-23 to form 1.24,25,25(OH)<sub>2</sub>D<sub>3</sub> mylated either at C-25 to form 1,24 25.28 (OH 4D. or at 76:te form 1 24.25.26(OH 4D). Thus, in an analogous thion, even though we do not have direct evidence of the weeking of 24.25 OH. Do not have direct evidence of the weeking of 24.25 OH. Do not both 24.25 of events.

25148- and Later, the trinydrolly itamin.

08(OH),  $O_2$  was not a significant circulating metabolite of min  $O_2$  in a vitamir  $C_2$  into verted rat as indicated by the ence of a radioactive peak in the migration position of an

increase a factories of pear in the migration position of an increase synthetic standard of 25.28(OH)<sub>2</sub>D<sub>2</sub> (Figure 5). In increase study at we did not have the synthetic standard 5.26(OH)<sub>2</sub>D<sub>2</sub>, we conclude the elution position 25.26(OH)<sub>2</sub>D<sub>2</sub> or our HPLC systems. As a result, we were

ble to conclude whether there was any formation of  $26(\mathrm{OH})_2\mathrm{D}_2$  in both our in v too and our in vivo studies.

ever, korzewsk, et al. 1989 in their recent study defi-lyestablished that 25,20(OH)<sub>2</sub>D<sub>2</sub> was not a major circu-

in or consider the part of the riamin D<sub>2</sub> intoxicated numbers the circulating level of itemit OH-D<sub>1</sub> can be as high as 250-750 ng/mL, and opter-come that develop in the distallination is being related the high circulating level of 21-OH-D<sub>1</sub> (Mawer et al., 1) Adm a recent story. As sewer, et al. (1988) invistigated metabolism of situmin D<sub>1</sub> in a stematic fashion in ny-diaminosis D<sub>2</sub>. They have indicated that both 24-OH-D<sub>2</sub> itemities in a vitamin antoxicated rat and that 14-OH-D<sub>2</sub> is mactivated through Donversion into 14-26(OH<sub>2</sub>D<sub>1</sub>). Until our present tudy, indirect indicated rational in available regarding the inactivation in way of 25-OH-D<sub>2</sub> is the conversion into 24-26(OH<sub>2</sub>D<sub>2</sub>) ones et al., 15-70 at 80. Our story further extending its one citization pathway of 25-OH-D<sub>2</sub> by demonstrating its one citization pathway of scenessing attemptive contribution into both 2-15-2-OH-D<sub>2</sub> by demonstrating its one citization pathway of scenessing attemptive contribution into both 2-15-2-OH-D<sub>2</sub> by demonstrating its one citization at the story of scenessing at a story of single-chain metabolism. dyrepresent significant ne attens i vitamin D<sub>1</sub> metatourm of the newly of solvered pathways of side-chain metablishment of the newly of solvered pathways of side-chain metablin out of the newly of solvered pathways of side-chain metablin out of the newly of solvered pathways of side-chain metablin out of the newly of solvered pathways of side-chain metablin out of the newly of solvered pathways of side-chain metablin out of the newly of solvered pathways of side-chain metablic out of the newly of solvered pathways of side-chain metabolic out of the newly of solvered pathways of side-chain metabolic out of the newly of solvered pathways of side-chain metabolic out of the newly of solvered pathways of side-chain metabolic out of the newly of solvered pathways of side-chain metabolic out of the newly of solvered pathways of side-chain metabolic out of the newly of solvered pathways of side-chain metabolic out of the newly of solvered pathways of side-chain metabolic out of the newly of solvered pathways of side-chain metabolic out of the newly of solvered pathways of side-chain metabolic out of the newly of solvered pathways of side-chain metabolic out of the newly of solvered pathways of side-chain metabolic out of the newly of side-chain metabolic out of the newly of side-chain metabolic out of si of the leading & Tserng, 1 (Kö). Studies to compare the biological pents.

It with of the two rest metabolites of 25-OH-De accombed in the previously we lettudied further metabolites of 25-OH-Ly in term of (a) calcium absorption by the of 25-OH-II; in term of (a) Lakinum absorption by the (b) calcium monifical on from the bone, and (a) their of (a)

binding affinity to the viturin D binding protein are presently in progress in our laboratory.

# ACKNOW LEDGMENTS

Wie gratefully acknowledge Dr. J. L. Napoli (State University of New York, Buffa.D. NY) for the kind gift of [3a-3H]vitamin Lig Dr. T. Kobayanni (Kobe Women's College of Pharmacy. Lobe, Japan) for the kind gift of authentic synthetic standard of 24 R),250 OH  $_2$ D $_2$ : Dr.  $_2$ P. L. Horr (USDA, Ames, IA+, Dr. M. R. Uskokav c (Hoffmann-La Fische Inc., Nutley, NJ), and Errs M. F. Holick and R. Ray (Botton University School of Medicine, Boston, MA) for many heipful discussions; F. Dayu for expert technical assistance and Lea Gold for efficient pecretarial assistance. This work is respectfully dedicated to the late Dr. E. G. Baggiothi (Hoffmann-La Roche, Inc. Nutle, NJ) without whose guidance. encouragement, and scientific advice the present work would have been impossible

Registry No. 23+OH+D $_1$ , 2-143-40-8, 24.23/OH  $_2$ D $_3$ , 75050+154-8, 24.0H-D-, 38(30-56-9) 24.21.25(OH),D-, 123942.85-8 24.25.28-(OH); D; 123997. 86.0

### Flerer Energy

Bligh, E. G. & Liver, W. J. (1989), Can. J. Biochem. Physiol. 1 37, 411-917.

Budgiblewicz, H. Djerass, C., & Wilham, D. F. (1967) Mais Specific metry of Organic Compounds, p. 03. Holcen-Day San Francisec

Horit, R. I., Feinhardt, T. A., Ramberg, C. F., K. szewski, N. J., & Nobell, J. L. (1986) J. Bicl. Chem. 261 925 -4216.

Johen, G., Schmess, H. K., & DeLuca, H. F. (1975) Biochemistry, 14, 2(4)-125%

Jones, G. Fosentha, A., Segev. D., Mazur, Y., Frolow, F., Halton Y. Rabinovich, D., & Snakken, Z. (1979) Blocnemistry 12, 1044-110

Jone . G., Schnier, H. F., Levan, L., & DeLuca, H. F. (1980) Arch Elochem Blomhys 202, 450-457

fores, G. Kans, E. Samaco, S., Furusaka, T., Takayama, H. & Suda, T. (1984) Elochemistry, 23, 1749-1754.

Jone, G., Vrierer, D. Lohner, D. Palda, V., & Edwards, N. S. 1-87) Stor ((154) 2043

Kerzew R. W. Esinnard, T. A., Napoli, J. L., Beitz, D. U. & Fiorit F. L. (1988) Bicchemistry 27, 5781-5790. Mawer E & Jiehr, J. T., Jucqueline, L. B., & Davies, M. (1933) Clin. Sc., 43, 135-14]

Napoli, I. L., Fizizini, M. A., Schnoes, H. E., & Felluca, H. F. (1979) Arch. Biochem. Biophys. 197. 119-125.

Norman A. W. Fott J., & Orol L. (1982) Endocr. Rev. 3. 33 -136.

Reda . G S v Tierng, E.-Y. (1986) Bid hemising 23. 3328-3386.

Reddy, G. S. Jones, C., Koch, S. W., & Frager, D. (1982) Am. J. Physial, 343, 8261-8271

Redoy, G. S., Jones, C., Koch, S. W., Fraser, C.I. & DeLuca. H. F. (1963) Am. J. Physic. 245, E359-E364.

Shepard, F., M., & DeLuca, H. F. (1980) Arch Blochem